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Description**BACKGROUND OF THE INVENTION**5 **Field of the Invention:**

[0001] The present invention relates to a coated cutting tool member that resists chipping and wear for long periods of time during cutting operations.

10 **Description of the Related Art**

[0002] Coated carbide cutting tool members are preferably composed of a tungsten carbide-based cemented carbide substrate and a hard coating layer preferably made of aluminum oxide (hereinafter referred to as "Al₂O₃"). Preferably, they further comprise a cubic-type titanium compound layer preferably including at least one layer of titanium compound having a "cubic" crystal structure preferably selected from titanium carbide (TiC), titanium nitride (TiN), titanium carbonitride (TiCN), titanium carboxide (TiCO), titanium nitroxide (TiNO) and titanium carbonitroxide (TiCNO). The hard coating layer is formed preferably by means of chemical vapor deposition and/or physical vapor deposition and have an average thickness of 3 to 20 µm. X-ray diffraction can confirm that the crystal structure of a titanium compound layer is cubic-type (hereinafter referred to as "cubic-type titanium compound layer"). A coated carbide cutting tool member having a hard coating layer, wherein the first layer is TiN, the second layer is TiCN, the third layer is TiCNO, the fourth layer is Al₂O₃ and fifth layer is TiN disclosed in Japanese Unexamined Patent Publication No.7-328810 (the contents of which are hereby incorporated by reference). These coated carbide cutting tool members are widely used in various fields of cutting operations, for example, continuous and interrupted cutting operation of metal work pieces.

[0003] It is known that cubic-type titanium compound layers have granular crystal morphology and are used for many applications. Recently, a TiCN layer that has a longitudinal crystal morphology has found use as highly wear resistant coating layer. TiC layers have been used as highly abrasion resistant materials in many applications. TiN layers have been used in many fields, for example, as an outermost layer of a coated cutting tool member and for various decorative products, because of its beautiful external view like gold. Layers of Al₂O₃ have several different crystal polymorphs, among which the alpha-Al₂O₃ is known as the thermodynamically most stable polymorph, having a corundum structure.

30 Typically, an Al₂O₃ coating formed by CVD has three kinds of Al₂O₃ polymorphs, namely, stable alpha-Al₂O₃, metastable kappa-Al₂O₃ and amorphous Al₂O₃.

[0004] In recent years, there has been an increasing demand for labor-saving, less time consuming cutting operations. These operations preferably include high speed cutting operations such as high speed feeding and/or high speed cutting. In these cutting operations, cutting tools are exposed to extraordinarily severe conditions. During these high speed cutting operations, the temperature of the cutting edge rises to 1000°C, or more and work chips of exceedingly high temperature are in contact with the surface of the rake face of the cutting tool. This phenomenon accelerates the occurrence of crater wear on the rake face. Thus, the cutting tool is chipped or damaged at a relatively early stage.

[0005] In order to circumvent this situation, a coated carbide cutting tool which has a relatively thick Al₂O₃ layer has been examined and produced. The Al₂O₃ layer has favorable properties such as extremely high resistance against oxidation, chemical stability and high hardness which meet the demands of cutting tools that are used under high temperature conditions. However, applying Al₂O₃ layers to cutting tools does not work out as one desires. Adhesion strength of the Al₂O₃ layer to an adjacent cubic-type titanium compound layer is usually not adequate, especially when the Al₂O₃ polymorph is alpha-type, and it is also inevitable that the Al₂O₃ layer has local nonuniformity in its thickness when it becomes a thicker layer. The Al₂O₃ layer tends to be thicker at the edge portion of the cutting tool, for example, than that at the other portions of the tool. When the thick Al₂O₃ layer is applied as a constituent of a hard coating layer, it is likely to show relatively short life time, for example, due to an occurrence of some kind of damage such as chipping, flaking and breakage.

[0006] As the cutting speed of various cutting operations continue to increase, thicker coatings of Al₂O₃ will be required to protect carbide cutting tools. With thicker Al₂O₃ layers, tool-life time will be more sensitive to both the adhesion strength between Al₂O₃ layer and cubic-type titanium compound layer as well as the toughness of Al₂O₃ layer itself. Methods for adhering Al₂O₃ layers to other compound layers and methods for making tough and thick Al₂O₃ layers continue to grow in importance with increasing demand for cutting tools that work at higher and higher speeds.

[0007] Cutting tool members are disclosed in EP-A-0 816 531 and US-A-4,463,062 which comprise a titanium oxide bonding layer with insufficient bonding to the Al₂O₃ layer.

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SUMMARY OF THE INVENTION

[0008] Accordingly, one object of this invention provides for a coated carbide cutting tool member having a thick

Al_2O_3 layer that strongly adheres to a cubic-type titanium compound layer and that shows excellent uniformity in Al_2O_3 thickness. Another object of the invention provides for coated carbide cutting tool members which have excellent wear resistance and damage resistance.

[0009] These and other objects of the present invention have been satisfied by the discovery of a coated carbide cutting tool member comprising a substrate and a hard coating layer on said substrate, wherein said hard coating layer comprises at least one layer comprising a titanium compound having a cubic lattice structure, at least one layer comprising aluminum oxide, and at least one intervening layer, wherein said intervening layer is between the layer comprising the titanium compound having a cubic lattice structure and the aluminum oxide layer, or between the aluminum oxide layers, and the intervening layer comprises titanium oxide having a corundum lattice structure (hereinafter referred to as " Ti_2O_3 ") and further comprises titanium carbonitride in a cubic lattice structure. This coated carbide cutting tool member gives good wear resistance and long tool lifetime when used in high speed cutting operations.

BRIEF DESCRIPTION OF THE DRAWING

[0010] A more complete appreciation of the invention and many of the attendant advantages thereof will be readily obtained as the same becomes better understood by reference to the following detailed description when considered in connection with the accompanying drawings, wherein:

Fig. 1 is a graph showing X-ray diffraction for coated carbide cutting inserts in accordance with the present invention in EXAMPLE 3, before the deposition of Al_2O_3 layer.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

[0011] The present invention provides for a cutting tool having a cutting tool member that is coated with a hard coating layer. A "cutting tool member" refers to the part of the cutting tool that actually cuts the work piece. Cutting tool members include exchangeable cutting inserts to be mounted on face milling cutter bodies, bit shanks of turning tools, and cutting blade of end mills. The cutting tool member is preferably made of tungsten carbide-based cemented carbide substrates.

[0012] A hard coating coats preferably a fraction of the surface, more preferably the entire surface of the cutting tool member. The hard coating comprises at least one layer comprising a titanium compound layer with a cubic lattice structure, at least one Al_2O_3 layer, and an intervening layer that lies between the titanium compound layer and the Al_2O_3 layer or between the Al_2O_3 layers. The intervening layer may directly contact one or both of the titanium compound layer with a cubic lattice structure and the Al_2O_3 layer. Although the Al_2O_3 layer is preferably the outermost layer of the hard coating layer, a TiN layer is used as outermost layer in many cases because of its beautiful appearance.

[0013] The titanium compound layer with the cubic lattice structure is composed of at least one layer selected from the group consisting of TiC, TiN, TiCN, TiCO, TiNO and TiCNO. The intervening layer comprises titanium oxide that has a corundum-type lattice structure (hereinafter referred to as " Ti_2O_3 ").

[0014] The preferred embodiments of the present invention were discovered after testing many different kinds of hard coating layers on coated carbide cutting tool members. In all of these tests, the hard coating layers included at least one titanium compound layer with a cubic lattice structure, at least one Al_2O_3 layer, and an intervening layer between the two other layers. From these tests, the following results (A) through (G) were found:

(A) When intervening layer preferably comprising Ti_2O_3 was inserted between said cubic-type titanium compound layer and said Al_2O_3 layer, the obtained coated carbide cutting tool exhibited longer tool life time.

(B) When intervening layer preferably comprising Ti_2O_3 was used, the cutting properties of the obtained cutting tool member varied according to the specific orientation in X-ray diffraction of said intervening layer. X-ray diffraction was performed using Cu $\text{K}\alpha$ -ray. When an intervening layer preferably comprises Ti_2O_3 having an X-ray diffraction pattern showing the maximum peak intensity at $2\theta = 53.8 \pm 1^\circ$ (the same as ASTM10-63, the entire contents of which are hereby incorporated by reference), the obtained coated carbide cutting tool member exhibited longer tool life time. Moreover, when intervening layer preferably comprises Ti_2O_3 having an X-ray diffraction pattern showing the maximum peak intensity at $2\theta = 34.5 \pm 1^\circ$, the obtained coated carbide cutting tool member exhibited an even longer lifetime.

(C) When an intervening layer preferably comprises Ti_2O_3 , having an X-ray diffraction pattern showing the maximum peak intensity at $2\theta = 34.5 \pm 1^\circ$, and further comprising a suitable amount of TiCNO, the obtained coated carbide cutting tool member exhibited even longer tool lifetimes in high speed continuous and interrupted cutting operations for steel and cast iron. The presence of TiCNO phase was confirmed by elemental analysis using an EPMA (electron probe micro analyzer) and X-ray diffraction. However, too much TiCNO in the intervening layer was not favorable because the properties of said layer became similar to that of cubic TiCNO layer.

(D) Other titanium oxide layers which can be obtained by chemical vapor deposition process including TiO , Ti_4O_7 and TiO_2 were also evaluated as intervening layers. The surface of these layers were smooth and dense nucleation

of Al_2O_3 was obtained for the intervening layers made from these materials just like for Ti_2O_3 . We thought that these phenomena might be attributed to the high density of oxygen atoms on the surface of said layers. For these layers, the presence of a cubic titanium compound phase was not confirmed. Coated carbide cutting inserts having intervening layers made from TiO , Ti_4O_7 and TiO_2 exhibited inferior cutting properties compared to the intervening layer comprising mainly Ti_2O_3 . Flaking of Al_2O_3 layer and chipping in quite early stages of cutting operation were frequently observed even in continuous cutting operations of steel and cast iron. For these observations we have found that Ti_2O_3 is the most preferred intervening layer between a cubic-type titanium compound layer and an Al_2O_3 layer.

(E) Improvement in cutting properties by having an intervening layer comprising mainly Ti_2O_3 might be attributed to the higher adhesion strength between this layer and the Al_2O_3 layer compared to the adhesion strength between a cubic-type titanium compound layer and an Al_2O_3 layer. We interpret the concept of "adhesion strength" as a combination effect of the "chemical bonding" between the two layers which are in contact with each other and the "mechanical bonding" between these two layers. An intervening layer preferably comprising Ti_2O_3 may have higher chemical bonding toward an Al_2O_3 layer than other cubic-type titanium compound layers and this layer may have more mechanical bonding because its surface is preferably rough. It has been confirmed that the surface morphology of the layer comprising mainly Ti_2O_3 is made favorably rougher, by the addition of a suitable amount of TiCNO in said layer. The positive effect of TiCNO in the layer comprising mainly Ti_2O_3 may be due to an increasing of mechanical bonding between said layer and the Al_2O_3 layer.

(F) The chemical bonding between other titanium oxide intervening layers, TiO , Ti_4O_7 and TiO_2 and the Al_2O_3 layer may also be high. However, the cutting properties of the coated carbide cutting tool member using these titanium oxides was found inadequate. We think that the reason for the relative short tool lifetime in cutting operations for these intervening layers might be attributed to a lack of a sufficient surface roughness. Consequently, the mechanical bonding between the intervening layers and the Al_2O_3 layer might have been weak.

(G) When the Al_2O_3 layer gets thicker, the tool lifetime of the coated carbide cutting tool member gets shorter. Experiments revealed that the shorter lifetime of the tool was caused by fracturing in the thick Al_2O_3 layer. The fracturing was attributed to a brittleness of thicker Al_2O_3 layers, especially at the edge of the tool member. This is because the Al_2O_3 layer at the edge is generally thicker than that at any other part of the tool, such as flank face or rake face.

30 [0015] In these cases, it is possible to make the thick Al_2O_3 layer tougher by replacing the thick Al_2O_3 with a composite structure layer preferably comprising at least two Al_2O_3 layers and at least one intervening layer preferably comprising mainly Ti_2O_3 . By this method, the nonuniformity in Al_2O_3 layer thickness was improved and consequently tool lifetime of said cutting tool member was improved even for an interrupted cutting operation.

35 [0016] Based on these results, the present invention provides for a coated carbide cutting tool member that exhibits extremely high wear resistance for various cutting operations and that has a long tool lifetime by providing a coated carbide cutting tool member preferably composed of a cemented carbide substrate and a hard coating layer preferably having an average thickness of 3 to 25 μm formed on said substrate being composed of at least one layer selected from the group of TiC , TiN , TiCN , TICO , TINO , TiCNO and Al_2O_3 , wherein said hard coating layer further has an intervening layer preferably comprising mainly Ti_2O_3 , having an X-ray diffraction pattern showing the maximum peak intensity at $2\theta = 34.5 \pm 1^\circ$, and formed between said cubic-type titanium compound layer and said Al_2O_3 layer. The present invention also provides for a coated carbide cutting tool member with a thick Al_2O_3 layer that exhibits extremely high toughness by providing a coated carbide cutting tool member, wherein the Al_2O_3 layer is replaced with a composite structure layer preferably comprising at least two Al_2O_3 layers and at least one intervening layer preferably comprising mainly Ti_2O_3 .

45 [0017] In the present invention, the average thickness of the hard coating layer is preferably 3 to 25 μm . Excellent wear resistance cannot be achieved at a thickness of less than 3 μm , whereas damage and chipping of the cutting tool member easily occur at a thickness of over 25 μm .

[0018] The average thickness of the intervening layer is preferably 0.1 to 5 μm . Satisfactory bonding effect toward both cubic-type titanium compound layer and Al_2O_3 layer cannot be achieved at a thickness of less than 0.1 μm , whereas the possibility of chipping occurrence of the cutting tool member becomes significant at a thickness of over 5 μm .

50 [0019] The average thickness of the individual Al_2O_3 layer in composite structure layer is preferably 0.5 to 12 μm , more preferably 0.5 to 10 μm , still more preferably 0.5 to 7 μm . It becomes difficult to provide satisfactory properties of Al_2O_3 such as oxidation resistance, chemical stability and hardness toward said composite structure layer at a thickness of less than 0.5 μm , whereas both the uniformity of layer thickness and toughness of said composite structure layer becomes insufficient at a thickness of over 12 μm .

55 [0020] The average thickness of the individual intervening layer in composite structure layer is preferably 0.05 to 2 μm . It becomes difficult to keep sufficient toughness of cutting tool member at a thickness of less than 0.05 μm , whereas

wear resistance decreases at a thickness of over 2 μm .

[0021] The ratio of TiCNO in an intervening layer comprising mainly Ti_2O_3 was expressed using ratio of carbon plus nitrogen in said layer as follows:

preferably

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$$0\% \leq (\text{C}+\text{N})/(\text{Ti}+\text{O}+\text{C}+\text{N}) \leq 10\%$$

more preferably

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$$0.5\% \leq (\text{C}+\text{N})/(\text{Ti}+\text{O}+\text{C}+\text{N}) \leq 5\%.$$

The properties of said layer were similar to that of a cubic TiCNO layer when the ratio was over 10%.

[0022] The "cubic" lattice structure is defined to include simple cubic lattices, body centered cubic lattices, and face centered cubic lattices, among others.

[0023] Further, said layer mainly comprising Ti_2O_3 is formed by means of chemical vapor deposition using a reactive gas preferably containing 0.4 to 10 percent by volume (hereinafter merely percent) of TiCl_4 , 0.4 to 10 percent of carbon dioxide (CO_2), 5 to 40 percent of nitrogen (N_2), 0 to 40 percent of argon (Ar), and the remaining balance of the reactive gas being hydrogen (H_2) at a temperature of 800 to 1100°C and a pressure of 30 to 500 Torr.

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EXAMPLES

[0024] Having generally described this invention, a further understanding can be obtained by reference to certain specific examples which are provided herein for purposes of illustration only and are not intended to be limiting unless otherwise specified.

EXAMPLE 1

[0025] The following powders were prepared as raw materials: a WC powder with an average grain size of 2.8 μm ; a coarse WC powder with an average grain size of 4.9 μm ; a TiC/WC powder with an average grain size of 1.5 μm (TiC/WC = 30/70 by weight); a (Ti,W)CN powder with an average grain size of 1.2 μm (TiC/TiN/WC = 24/20/56); a TaC/NbC powder with an average grain size of 1.2 μm (TaC/NbC = 90/10); and a Co powder with an average grain size of 1.1 μm . These powders were compounded based on the formulation shown in Table 1, wet-mixed in a ball mill for 72 hours, and dried. The dry mixture was pressed to form a green compact for cutting insert defined in ISO-CNMG120408 (for carbide substrates A through D) or ISO-SEEN42AFTN1 (for carbide substrate E), followed by vacuum sintering under the conditions set forth in Table 1 for Carbide substrates A through E. (Note: the contents of ISO-CNMG120408 and ISO-SEEN42AFTN1 are hereby incorporated by reference.)

[0026] The carbide substrate B was held in a CH_4 atmosphere of 100 Torr at 1400°C for 1 hour, followed by annealing for carburization. The carburized substrate was then subjected to treatment by acid and barrel finishing to remove carbon and cobalt on the substrate surface. The substrate was covered with a Co-enriched zone having a thickness of 42 μm and a maximum Co content of 15.9 percent by weight at a depth of 11 μm from the surface of the substrate.

[0027] Sintered carbide substrates A and D had a Co-enriched zone having a thickness of 23 μm and a maximum Co content of 9.1 percent by weight at a depth of 17 μm from the surface of the substrate. Carbide substrates C and E had no Co-enriched zone and had homogeneous microstructures.

[0028] The Rockwell hardness (Scale A) of each of the carbide substrates A through E is also shown in Table 1.

[0029] The surface of the carbide substrates A through E were subjected to honing and chemical vapor deposition using conventional equipment under the conditions shown in Table 2 to form hard coating layers that had a composition and a designed thickness (at the flank face of the cutting insert) shown in Tables 3 and 4. TiCN* in each Table represented the TiCN layer that had a crystal morphology longitudinally grown as described in Japanese Unexamined Patent Publication No-6-8010 (the entire contents of which are hereby incorporated by reference). Coated carbide cutting inserts in accordance with the present invention 1 through 10 and conventional coated carbide cutting inserts 1 through 10 were produced in such a manner.

[0030] Further, continuous cutting tests and interrupted cutting tests were conducted for above cutting inserts under the following conditions.

[0031] A wear width on a flank face was measured in each tests.

[0032] For coated carbide cutting inserts of the present invention 1 through 9 and conventional coated carbide cutting inserts 1 through 9, the following cutting tests were conducted:

(1-1) Cutting style: Continuous turning of alloy steel
 Work piece: JIS SCM440 round bar
 Cutting speed: 350 m/min
 Feed rate: 0.4 mm/rev
 5 Depth of cut: 3 mm
 Cutting time: 10 min
 Coolant: Dry

(1-2) Cutting style: Interrupted turning of alloy steel
 10 Work piece: JIS SNCM439 square bar
 Cutting speed: 180 m/min.
 Feed rate: 0.25 mm/rev.
 Depth of cut: 3 mm
 Cutting time: 5 min Coolant: Dry

15 [0033] For coated carbide cutting inserts of the present invention 10 and conventional coated carbide cutting inserts 10, following cutting tests were conducted:

(1-3) Cutting style: Milling of carbon steel
 20 Work piece: JIS S45C square bar (100 mm width x 500 mm length)
 Cutting tool configuration: single cutting insert mounted with a cutter of 125 mm diameter
 Cutting speed: 200 m/min
 Feed rate: 0.15 mm/tooth
 25 Depth of cut: 2 mm
 Cutting time: 10 min
 Coolant: Dry

Results were shown in Table 5.

30 EXAMPLE 2

[0034] The same carbide substrates A through E as in EXAMPLE 1 were prepared. The surfaces of the carbide substrates A through E were subjected to honing and chemical vapor deposition using conventional equipment under the conditions shown in Table 6 to form hard coating layers that had a composition and a designed thickness (at the 35 flank of the cutting insert) shown in Table 7 and 8. Coated carbide cutting inserts in accordance with the present invention 11 through 20 and conventional coated carbide cutting inserts 11 through 20 were produced in such a manner.

[0035] Further, continuous cutting tests and interrupted cutting tests were conducted for above cutting inserts under the following conditions. A wear width on a flank face was measured in each test.

[0036] For coated carbide cutting inserts of the present invention 11, 12 and conventional coated carbide cutting 40 inserts 11, 12, following cutting tests were conducted:

(2-1) Cutting style: Intenupted turning of Ductile cast iron
 Work piece: JIS FCD450 square bar
 Cutting speed: 250 m/min
 45 Feed rate: 0.25 mm/rev
 Depth of cut: 2 mm
 Cutting time: 5 min
 Coolant: Dry

50 [0037] For coated carbide cutting inserts of the present invention 13, 14 and conventional coated carbide cutting inserts 13, 14, following cutting tests were conducted:

(2-2) Cutting style: Interrupted turning of Alloy steel
 Work piece: JIS SCM415 square bar
 55 Cutting speed: 250 m/min
 Feed rate: 0.25 mm/rev
 Depth of cut: 2 mm
 Cutting time: 5 min

Coolant: Dry

[0038] For coated carbide cutting inserts of the present invention 15, 16 and conventional coated carbide cutting inserts 15, 16, following cutting tests were conducted:

5 (2-3) Cutting style: Interrupted turning of Carbon steel
Work piece: JIS S45C square bar
Cutting speed: 250 m/min
Feed rate: 0.25 mm/rev
10 Depth of cut: 2 mm
Cutting time: 5 min Coolant: Dry

[0039] For coated carbide cutting inserts of the present invention 17, 18 and conventional coated carbide cutting inserts 17, 18, following cutting tests were conducted:

15 (2-4) Cutting style: Interrupted turning of Cast iron
Work piece: JIS FC200 square bar
Cutting speed: 250 m/min
Feed rate: 0.25 mm/rev
20 Depth of cut: 2 mm
Cutting time: 5 min Coolant: Dry

[0040] For coated carbide cutting inserts of the present invention 19, 20 and conventional coated carbide cutting inserts 19, 20, following cutting tests were conducted:

25 (2-5) Cutting style: Milling of Alloy steel
Work piece: JIS SCM440 square bar (100 mm width x 500 mm length)
Cutting tool configuration: single cutting insert mounted with a cutter of 125 mm diameter
Cutting speed: 250 m/min
30 Feed rate: 0.2 mm/tooth
Depth of cut: 2 mm
Cutting time: 8.6 min
Coolant: Dry

35 Results were shown in Table 9.

EXAMPLE 3

[0041] The same carbide substrate A as in EXAMPLE 1 was prepared. The surfaces of the carbide substrate A were subjected to honing and chemical vapor deposition using conventional equipment under the conditions shown in Table 10 to form hard coating layers that had a composition and a designed thickness (at the flank of the cutting insert) shown in Table 11. Coated carbide cutting inserts in accordance with the present invention 21 through 29 and conventional coated carbide cutting insert 21 were produced in such a manner.

[0042] Intervening layers comprising mainly Ti_2O_3 of the cutting inserts of present invention 21 through 29 and a cubic-type TiCNO layer of the cutting insert of conventional invention 21 were subjected to elemental analysis using an EPMA (electron probe micro analyzer) or ABS (auger electron spectroscopy). The cutting insert used in the elemental analysis was identical to the one used in the cutting test. The elemental analysis was carried out by irradiating an electron beam having a diameter of 1 μm onto the center of the flank face. These layers were also subjected to X-ray diffraction analysis using Cu K α -ray. Analytical results using a ratio of carbon plus nitrogen in each layer, $(C + N) / (Ti + O + C + N)$, were shown in Table 12.

[0043] Further, continuous cutting tests were conducted for above cutting inserts under the following conditions: A wear width on a flank face was measured in each tests.

[0044] For coated carbide cutting inserts of the present invention 21 through 29 and conventional coated carbide cutting insert 21, following cutting tests were conducted:

55 (3-1) Cutting style: Continuous turning of alloy steel
Work piece: JIS SNCM439 round bar
Cutting speed: 280 m/min

Feed rate: 0.35 mm/rev
 Depth of cut: 1.0 mm
 Cutting time: 10 min Coolant: Dry

5 Results were shown in Table 12.

EXAMPLE 4

[0045] The same carbide substrate A as in EXAMPLE 1 was prepared. The surface of the carbide substrate A was subjected to honing and chemical vapor deposition using conventional equipment under the conditions shown in Table 13 to form hard coating layers that had a composition and a designed thickness (at the flank of the cutting insert) shown in Table 14. Coated carbide cutting inserts in accordance with the present invention 30 through 34 and conventional coated carbide cutting inserts 22 through 26 were produced in such a manner.

[0046] Further, continuous cutting tests and interrupted cutting tests were conducted for the above cutting inserts under the following conditions. A wear width on a flank face was measured in each tests.

[0047] For coated carbide cutting inserts of the present invention 30 through 34 and conventional coated carbide cutting inserts 22 through 26, following cutting tests were conducted:

(4-1) Cutting style: Continuous turning of carbon steel
 Work piece: JIS S45C round bar
 Cutting speed: 450 m/min
 Feed rate: 0.3 mm/rev
 Depth of cut: 3 mm
 Cutting time: 10 min Coolant: Dry

(4-2) Cutting style: Interrupted turning of carbon steel
 Work piece: JIS S45C square bar
 Cutting speed: 200 m/min
 Feed rate: 0.3 mm/rev
 Depth of cut: 3 mm
 Cutting time: 5 min
 Coolant: Dry

Results were shown in Table 15.

EXAMPLE 5

[0048] A cemented carbide cutting tool member of the present invention is coated with the following series of layers to form a hard coating layer:

6th layer	TiN	0.3 microns thick
5th layer	Al ₂ O ₃	3 microns thick
4th layer	TiC	1 micron thick
3rd layer	Al ₂ O ₃	10 microns thick
2nd layer	Mostly Ti ₂ O ₃	1 micron thick
1st layer	TiCN	5 microns thick
Substrate	Cemented Carbide	

50 [0049] The present application is based on Japanese Priority Applications JP 09-120704, filed on May 12, 1997, JP 09-238198, filed on September 3, 1997, and JP 09-318100, filed on November 19, 1997, the entire contents of which are hereby incorporated by reference.

Table 1

Carbide substrate	Composition (wt%)					Vacuum sintering conditions			Rockwell hardness (Scale A) (HRA)
	Co	(Ti,W) C	(Ti,W)CN	(Ta,Nb)C	WC	Vacuum (torr)	Temperature (°C)	Time (hr)	
A	6.3	—	6	4.1	Balance	0.10	1380	1	90.3
B	5.3	5.2	—	5.1	Balance	0.05	1450	1	90.9
C	9.5	8.1	—	4.9	Balance	0.05	1380	1.5	89.9
D	4.5	—	4.8	3.1	Balance	0.10	1410	1	91.4
E	10.2	—	—	2.2	Balance (Coarse)	0.05	1380	1	89.7

Table 2

Hard coating layer	Conditions for forming hard coating layer		
	Composition of reactive gas (volume %)	Pressure (torr)	Temperature (°C)
Al2O3	AlCl3 : 2.2% , CO2 : 5.5% , HCl : 2.2% , H2 : Balance	50	1000
TiC	TiCl4 : 4.2% , CH4 : 4.5% , H2 : Balance	50	1020
TiN	TiCl4 : 4.2% , N2 : 30% , H2 : Balance	200	1020
TiCN	TiCl4 : 4.2% , CH4 : 4% , N2 : 20% , H2 : Balance	50	1020
TiCN*	TiCl4 : 4.2% , CH3CN : 0.6% , N2 : 20% , H2 : Balance	50	910
TiCO	TiCl4 : 2% , CO : 6% , H2 : Balance	50	980
TiNO	TiCl4 : 2% , NO : 6% , H2 : Balance	50	980
TiCNO	TiCl4 : 2% , CO : 3% , N2 : 30% , H2 : Balance	50	980
Ti2O3**	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 43.5% , H2 : Balance	200	1000

* : TiCN layer having a crystal morphology longitudinally grown
 ** : intervening layer comprising mainly corundum titanium oxide

Table 3

Insert	Substrate	Hard coating layer (Figure in parentheses means designed thickness ; μ m)					
		First layer	Second layer	Third layer	Fourth layer	Fifth layer	Sixth layer
This invention	1 A	TIN (0.1)	TiCN* (5)	Ti2O3** (0.1)	Al2O3 (3)	TiN (0.2)	
	2 B	TiC (0.5)	TiN (1)	TiCN* (4)	Ti2O3** (1.5)	Al2O3 (4)	
	3 C	TIN (0.1)	TiCN* (3)	TiCO (0.1)	Ti2O3** (2.5)	Al2O3 (4)	TiN (0.1)
	4 D	TIN (0.1)	TiCN* (3)	TiCNO (0.1)	Ti2O3** (0.5)	Al2O3 (4.5)	
	5 A	TiCN (3)	TiCN* (6)	TiN (2.5)	Ti2O3** (4.5)	Al2O3 (2)	
	6 B	TiC (1)	TiCN* (5)	TiNO (0.1)	TiCNO (0.3)	Ti2O3** (1.5)	Al2O3 (4)
	7 C	TIN (0.5)	TiCN (5)	Ti2O3** (0.3)	Al2O3 (4)	Ti2O3** (1)	TiN (0.3)
	8 D	TiC (3)	Ti2O3** (5)	Al2O3 (2)			
	9 A	TIN (1)	Ti2O3** (1)	Al2O3 (3)	Ti2O3** (1)	TiCN* (4)	Al2O3 (4)
	10 B	TIN (0.1)	TiCN* (5)	TiC (3)	TiNO (0.1)	Ti2O3** (1)	Al2O3 (3)

* : TiCN layer having a crystal morphology longitudinally grown
** : intervening layer comprising mainly corundum titanium oxide

Table 4

Insert	Substrate	Hard coating layers (Figure in parentheses means designed thickness ; μ m)				
		First layer	Second layer	Third layer	Fourth layer	Fifth layer
Conventional	1 A	TiN (0.1)	TiCN* (5)	Al2O3 (3)	TiN (0.2)	
	2 B	TiC (0.5)	TiN (1)	TiCN* (4)	Al2O3 (4)	
	3 C	TiN (0.1)	TiCN* (3)	TiCO (0.1)	Al2O3 (4)	TiN (0.1)
	4 D	TiN (0.1)	TiCN* (3)	TiCNO (0.1)	Al2O3 (4.5)	
	5 A	TiCN (3)	TiCN* (6)	TiN (2.5)	Al2O3 (2)	
	6 B	TiC (1)	TiCN* (5)	TiNO (0.1)	TiCNO (0.1)	Al2O3 (4)
	7 C	TiN (0.5)	TiCN (5)	Al2O3 (4)	TiN (0.3)	
	8 D	TiC (3)	Al2O3 (2)			
	9 A	TiN (1)	Al2O3 (3)	TiCN* (4)	Al2O3 (4)	
	10 E	TiN (0.1)	TiCN* (5)	TiC (3)	TiNO (0.1)	Al2O3 (3)
						TiN (0.1)

* : TiCN layer having a crystal morphology longitudinally grown

Table 5

Insert	Flank wear (mm)			Insert	Flank wear (mm)		
	(1-1)	(1-2)	(1-3)		(1-1)	(1-2)	(1-3)
This invention	1 0.25	0.19	—	1	0.29	Failure at 1.0min	—
	2 0.22	0.17	—	2	0.28	Failure at 0.5min	—
	3 0.30	0.18	—	3	0.30	Failure at 3.5min	—
	4 0.24	0.19	—	4	0.25	Failure at 1.0min	—
	5 0.29	0.20	—	5	0.29	Failure at 1.0min	—
	6 0.21	0.20	—	6	0.25	Failure at 1.0min	—
	7 0.29	0.21	—	7	0.31	Failure at 0.5min	—
	8 0.21	0.20	—	8	0.24	Failure at 0.5min	—
	9 0.22	0.18	—	9	0.30	Failure at 2.0min	—
	10 —	—	0.20	10	—	—	Failure at 4.0min

Remark : Failure is caused by chipping

Table 6

Hard coating layer	Conditions for forming hard coating layer		
	Composition of reactive gas (volume %)	Pressure (torr)	Ambience (°C)
TiC	TiCl4 : 4% , CH4: 9% , H2 : Balance	50	1020
TiN (first layer)	TiCl4 : 4% , N2 : 30% , H2 : Balance	50	920
TiN (the other layer)	TiCl4 : 4% , N2 : 35% , H2 : Balance	200	1020
TiCN*	TiCl4 : 4% , CH3CN : 1.2% , N2: 30% , H2 : Balance	50	900
TiCN	TiCl4 : 4% , CH4: 4% , N2 : 30% , H2 : Balance	50	1020
TiCO	TiCl4 : 4% , CO : 9% , H2 : Balance	50	1020
TiNO	TiCl4 : 4% , NO : 9% , H2 : Balance	50	1020
TiCNO	TiCl4 : 4% , CO : 5% , N2 : 8% , H2 : Balance	50	1020
Ti2O3**	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 43.5% , H2 : Balance	80	1020
Al2O3 (a)	AlCl3 : 2.2% , CO2 : 5.5% , HCl : 2.2% , H2 : Balance	50	1030
Al2O3 (b)	AlCl3 : 2.2% , CO2 : 5.5% , HCl : 2.2% , H2 : Balance	50	970

* : TiCN layer having a crystal morphology longitudinally grown

** : intervening layer comprising mainly corundum titanium oxide

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Table 7

Insert	Substrate	Hard coating layer (Figure in parentheses means designed thickness ; μ m)			Composite layer			Outermost layer
		First layer	Second layer	Third layer	Initial layer	Medium layer	Final layer	
11	A	TiN (0.5)	TiCN* (9)	TiCO (0.3)	Al2O3 (b) (1.5)	Ti2O3** (0.1)	Al2O3 (b) (1.5)	TiN (0.5)
12	A	TiN (0.5)	TiCN* (5)	TiCN (2)	Al2O3 (a) (1)	Ti2O3** (0.1) : 3 layers Al2O3 (a) (2) : 2 layers	Al2O3 (a) (1)	—
13	B	TiC (2)	TiCO (1)	TiCN* (5)	Al2O3 (a) (1)	Ti2O3** (0.2) : 10 layers Al2O3 (a) (2) : 9 layers	Al2O3 (a) (1)	TiN (0.5)
14	B	TiC (5)	—	—	Al2O3 (a) (1)	Ti2O3** (0.1) : 3 layers Al2O3 (a) (1.5) : 2 layers	Al2O3 (a) (1)	—
15	C	TiCN (3)	TiCN* (3)	TiCN0 (0.5)	Al2O3 (b) (2)	Ti2O3** (0.3) : 2 layers Al2O3 (b) (2) : 1 layer	Al2O3 (b) (2)	—
This invention	C	TiN (2)	TiCN* (5)	—	Al2O3 (a) (3)	Ti2O3** (0.2) : 2 layers Al2O3 (b) (3) : 1 layer	Al2O3 (a) (3)	—
	D	TiN (1)	TiCN* (5)	TiCO (0.3)	Al2O3 (a) (7)	Ti2O3** (0.3)	Al2O3 (a) (7)	TiN (1)
18	D	TiN (1)	TiCN* (5)	TiNO (0.5)	Al2O3 (a) (0.5)	Ti2O3** (0.1) : 15 layers Al2O3 (a) (1) : 14 layer	Al2O3 (a) (0.5)	TiN (1)
19	E	TiC (2)	—	—	Al2O3 (a) (0.5)	Ti2O3** (0.05) : 2 layers Al2O3 (a) (0.5) : 1 layer	Al2O3 (a) (0.5)	—
20	B	TiCN (3)	—	—	Al2O3 (b) (1.5)	Ti2O3** (0.1)	Al2O3 (b) (1.5)	—

*: TiCN layer having a crystal morphology longitudinally grown

**: intervening layer comprising mainly corundum titanium oxide

Table 8

Insert	Substrate	Hard coating layer (Figure in parentheses means designed thickness ; μ m)				
		First layer	Second layer	Third layer	Fourth layer	Fifth layer
Conventional	11 A	TiN (0.5)	TiCN* (9)	TiCO (0.3)	Al2O3 (b) (3)	TiN (0.5)
	12 A	TiN (0.5)	TiCN* (5)	TiCN (2)	Al2O3 (a) (6)	—
	13 B	TiC (2)	TiCO (1)	TiCN* (5)	Al2O3 (a) (5)	TiN (0.5)
	14 B	TiC (5)	Al2O3 (a) (7)	—	—	—
	15 C	TiCN (3)	TiCN* (3)	TiCNO (0.5)	Al2O3 (b) (8)	—
	16 C	TiN (2)	TiCN* (5)	Al2O3 (a) (2)	TiN (1)	—
	17 D	TiN (1)	TiCN* (5)	TiCO (0.3)	Al2O3 (a) (14)	TiN (1)
	18 D	TiN (1)	TiCN* (5)	TiNO (0.5)	Al2O3 (a) (15)	TiN (1)
	19 E	TiC (2)	Al2O3 (a) (2)	—	—	—
	20 E	TiCN (3)	Al2O3 (b) (3)	TiN (0.3)	—	—

* : TiCN layer having a crystal morphology longitudinally grown

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Table 9

Insert	Flank wear (mm)	Insert	Flank wear (mm)
11	0.17	Conventional	11 Failure at 0.9min
12	0.18		12 Failure at 1.4min
13	0.21		13 Failure at 2.1min
14	0.20		14 Failure at 2.5min
15	0.18		15 Failure at 1.1min
16	0.18		16 Failure at 2.3min
17	0.17		17 Failure at 2.5min
18	0.15		18 Failure at 1.6min
19	0.21		19 Failure at 3.3min
20	0.22		20 Failure at 1.6min

Remark : Failure is caused by chipping

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Table 10

Hard coating layer	Composition of reactive gas (volume %)	Conditions for forming hard coating layer	
		Pressure (torr)	Ambience
TiN (first layer)	TiCl4 : 4% , N2 : 30% , H2 : Balance	50	920
TiN (the other layer)	TiCl4 : 4% , N2 : 35% , H2 : Balance	200	1020
TiCN*	TiCl4 : 4% , CH3CN : 1.2% , N2 : 30% , H2 : Balance	50	900
TiCNO	TiCl4 : 4% , CO : 5% , N2 : 8% , H2 : Balance	50	1020
Ti2O3** (a)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 30% , Ar : 40% , H2 : Balance	200	1030
Ti2O3** (b)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 20% , Ar : 30% , H2 : Balance	200	1030
Ti2O3** (c)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 20% , Ar : 20% , H2 : Balance	200	1030
Ti2O3** (d)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 20% , Ar : 10% , H2 : Balance	200	1030
Ti2O3** (e)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 10% , Ar : 5% , H2 : Balance	200	1030
Ti2O3** (f)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 10% , Ar : 0% , H2 : Balance	200	1030
Ti2O3** (g)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 10% , Ar : 5% , H2 : Balance	50	900
Ti2O3** (h)	TiCl4 : 2.5% , CO2 : 3.5% , N2 : 5% , Ar : 5% , H2 : Balance	100	950
Ti2O3** (i)	TiCl4 : 2.5% , CO2 : 2.0% , N2 : 5% , Ar : 0% , H2 : Balance	250	1030
Al2O3	AlCl3 : 2.2% , CO2 : 5.5% , HCl : 2.2% , H2 : Balance	50	1030

*: TiCN layer having a crystal morphology longitudinally grown

**: intervening layer comprising mainly corundum titanium oxide

Table 11

Insert	Hard coating layer (Figure in parentheses means designed thickness : μ m)					
	1st layer	2nd layer	3rd layer	4th layer	5th layer	
21	TiN (1)	TiCN* (6)	Ti2O3** (a) (1)	Al2O3 (7)	TiN (0.3)	
22	TiN (1)	TiCN* (6)	Ti2O3** (b) (1)	Al2O3 (7)	TiN (0.3)	
23	TiN (1)	TiCN* (6)	Ti2O3** (c) (1)	Al2O3 (7)	TiN (0.3)	
24	TiN (1)	TiCN* (6)	Ti2O3** (d) (1)	Al2O3 (7)	TiN (0.3)	
This invention	25	TiN (1)	TiCN* (6)	Ti2O3** (e) (1)	Al2O3 (7)	TiN (0.3)
	26	TiN (1)	TiCN* (6)	Ti2O3** (f) (1)	Al2O3 (7)	TiN (0.3)
	27	TiN (1)	TiCN* (6)	Ti2O3** (g) (1)	Al2O3 (7)	TiN (0.3)
	28	TiN (1)	TiCN* (6)	Ti2O3** (h) (1)	Al2O3 (7)	TiN (0.3)
	29	TiN (1)	TiCN* (6)	Ti2O3** (i) (1)	Al2O3 (7)	TiN (0.3)
	Conventional	21	TiN (1)	TiCN* (6)	TiCNO (1)	Al2O3 (7)

* : TiCN layer having a crystal morphology longitudinally grown
 ** : intervening layer comprising mainly corundum titanium oxide

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Table.12

Insert	$(C+N) / (Ti+O+C+N)$	Analytical data		Flank wear (mm)
		Position of maximum peak in XRD pattern of Ti_2O_3 layer	$2\theta = 34.5^\circ$	
21	0%		$2\theta = 34.5^\circ$	0.43
22	0.7%		$2\theta = 34.5^\circ$	0.29
23	2.4%		$2\theta = 34.5^\circ$	0.24
24	4.6%		$2\theta = 34.5^\circ$	0.31
25	8.1%		$2\theta = 34.5^\circ$	0.38
This invention			$2\theta = 34.5^\circ$	
26	14.1%		$2\theta = 34.5^\circ$	0.42
27	1.8%		$2\theta = 54.0^\circ$	0.40
28	3.2%		$2\theta = 24.1^\circ$	0.44
29	17.6%		$2\theta = 54.0^\circ$	0.50
Conventional	32.2%		—	0.68

Table 13

Hard coating layer	Conditions for forming hard coating layer		
	Composition of reactive gas (volume %)	Pressure (torr)	Temperature (°C)
TiC	TiCl ₄ : 4% , CH ₄ : 9% , H ₂ : Balance	50	1020
TiN	TiCl ₄ : 4% , N ₂ : 35% , H ₂ : Balance	200	1020
TiCN	TiCl ₄ : 4% , CH ₄ : 4% , N ₂ : 30% , H ₂ : Balance	50	1020
TiCN*	TiCl ₄ : 4% , CH ₃ CN : 1.2% , N ₂ : 30% , H ₂ : Balance	50	900
TiCO	TiCl ₄ : 4% , CO : 4% , H ₂ : Balance	50	1020
TiNO	TiCl ₄ : 4% , NO : 6% , H ₂ : Balance	50	1020
TiCNO	TiCl ₄ : 4% , CO : 3% , N ₂ : 30% , H ₂ : Balance	50	1020
Ti ₂ O ₃ **	TiCl ₄ : 3% , CO ₂ : 3% , N ₂ : 30% , H ₂ : Balance	100	1020
Al ₂ O ₃	AlCl ₃ : 2.2% , CO ₂ : 5.5% , HCl : 2.2% , H ₂ : Balance	50	1020

* : TiCN layer having a crystal morphology longitudinally grown

** : intervening layer comprising mainly corundum titanium oxide

Table 14

Insert	Hard coating layer (Figure in parentheses means designed thickness; μ m)									
	First layer	Second layer	Third layer	Fourth layer	Fifth layer	Sixth layer	Seventh layer	Eighth layer	Ninth layer	Tenth layer
This invention	30 TiN (0.5) TiCN* (6)	Ti2O3** (0.8)	Al2O3 (5)	Ti2O3** (0.2)	Al2O3 (4)	TiN (0.3)				
	31 TiN (0.3) TiCN* (5)	TiC (3)	Ti2O3** (0.5)	Al2O3 (4)	Ti2O3** (0.2)	Al2O3 (4)	TiN (0.3)			
	32 TiCN (5) Ti2O3** (0.5)	Al2O3 (4)	Ti2O3** (0.1)	Al2O3 (3)	Ti2O3** (0.1)	Al2O3 (3)	TiN (0.3)			
	33 TiC (6) Ti2O3** (0.8)	Al2O3 (5)	Ti2O3** (0.2)	Al2O3 (5)	Ti2O3** (0.2)	Al2O3 (5)	TiN (0.3)			
	34 TiN (0.5) TiCN* (5)	Ti2O3** (0.5)	Al2O3 (3)	Ti2O3** (0.2)	Al2O3 (3)	Ti2O3** (0.2)	Al2O3 (3)	TiN (0.3)		
	22 TiN (0.5) TiCN* (6)	TiCN (0.4)	Al2O3 (9)	TiN (0.3)						
Conventional	23 TiN (0.3) TiCN* (5)	TiC (3)	TiN (0.5)	Al2O3 (12)	TiN (0.3)					
	24 TiCN (5) TiCO (0.5)	Al2O3 (10)								
	25 TiC (6) TiNO (0.4)	Al2O3 (15)	TiN (0.3)							
	26 TiN (0.5) TiCN* (5)	TiCO (0.4)	Al2O3 (3)	TiN (0.2)	Al2O3 (3)	TiN (0.2)	Al2O3 (3)	TiN (0.3)		

* : TiCN layer having a crystal morphology longitudinally grown
 ** : intervening layer comprising mainly corundum titanium oxide

Table 15

Insert	Flank wear (mm)		Insert	Flank wear (mm)	
	(4-1)	(4-2)		(4-1)	(4-2)
30	0.31	0.25		22	0.36
31	0.32	0.24		23	0.33
This invention	0.29	0.28	Conventional	24	0.49
33	0.30	0.25		25	0.57
34	0.33	0.24		26	0.33

Remark : Failure is caused by chipping

Claims

1. A coated carbide cutting tool member comprising;

5 a substrate and
 a hard coating layer on said substrate,
 wherein said hard coating layer comprises at least one layer comprising a titanium compound having a cubic lattice structure, at least one layer comprising aluminum oxide, and at least one intervening layer,
 wherein said intervening layer is between said layer comprising said titanium compound having a cubic lattice structure and said aluminum oxide layer, or between said aluminum oxide layers, and
 10 said intervening layer comprises titanium oxide having a corundum lattice structure and further comprises titanium carbonitroxide in a cubic lattice structure.

2. The article of Claim 1, wherein said substrate comprises tungsten carbide,

15 3. The article of Claim 1 or 2, wherein said at least one layer comprising said titanium compound having a cubic lattice structure comprises at least one layer selected from the group consisting of titanium carbide, titanium nitride, titanium carbonitride, titanium carboxide, titanium nitroxide and titanium carbonitroxide.

20 4. The article of any of Claims 1 to 3, wherein said intervening layer has a thickness of 0.1 to 5 μm .

5 5. The article of any of Claims 1 to 3, wherein said intervening layer has a thickness of 0.05 to 2 μm .

6. The article of any of Claims 1 to 5, wherein said hard coating layer has a thickness of 3 to 25 μm .

25 7. The article of any of Claims 1 to 6, wherein each of said aluminum oxide layers has a thickness of 0.5 to 10 μm .

8. The article according to any of Claims 1 to 7, wherein said intervening layer comprising titanium oxide having a corundum lattice structure shows a maximum peak intensity at $2\theta = 34.5 \pm 1^\circ$ in a X-ray diffraction pattern using a Cu $\text{K}\alpha$ -ray.

30 9. The article according to any of Claims 1 to 8, wherein the atomic ratio of carbon, nitrogen, oxygen and titanium in said intervening layer is expressed as follows:

35 $0\% \leq (\text{C}+\text{N})/(\text{Ti} + \text{O} + \text{C} + \text{N}) \leq 10\%$.

10. The article according to Claim 9, wherein said atomic ratio is:

40 $0.5\% \leq (\text{C} + \text{N}) / (\text{Ti} + \text{O} + \text{C} + \text{N}) \leq 5\%$.

11. A coated carbide cutting tool member comprising;

45 a substrate comprising tungsten carbide and
 a hard coating layer on said substrate having a thickness of 3 to 25 μm ,
 wherein said hard coating layer comprises at least one layer comprising a titanium compound having a cubic lattice structure, at least two layers comprising aluminum oxide, and at least one intervening layer,
 wherein said intervening layer is between said layer comprising said titanium compound having a cubic lattice structure and said aluminum oxide layer or between said aluminum oxide layers, and
 50 said intervening layer comprises titanium oxide having a corundum lattice structure and further comprises titanium carbonitroxide in a cubic lattice structure.

55 12. The article of Claim 11, wherein said at least one layer comprising said titanium compound having a cubic lattice structure comprises at least one layer selected from the group consisting of titanium carbide, titanium nitride, titanium carbonitride, titanium carboxide, titanium nitroxide and titanium carbonitroxide.

13. The article according to Claim 11 or 12, wherein each of said aluminum oxide layers has a thickness of 0.5 to 10 μm .

14. The article of any of Claims 11 to 13, wherein said intervening layer has a thickness of 0.05 to 2 μm .

15. The article according to any of Claims 11 to 14, wherein said intervening layer comprising titanium oxide having a corundum lattice structure shows a maximum peak intensity at $2\theta = 34.5 \pm 1^\circ$ in a X-ray diffraction pattern using a Cu $\text{k}\alpha$ -ray.

5 16. The article according to any of Claims 1 to 15, wherein said intervening layer is in contact with both of said layer comprising said titanium compound having a cubic lattice structure, and said aluminum oxide layer.

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Patentansprüche

1. Beschichtetes Carbid-Schneidewerkzeugelement umfassend:

15 ein Substrat und
auf dem Substrat eine harte Deckschicht
worin die harte Deckschicht mindestens eine Schicht, die eine Titanverbindung mit kubischer Gitterstruktur
enthält, mindestens eine Schicht, die Aluminiumoxid enthält und mindestens eine Zwischenschicht umfasst,
wobei die Zwischenschicht zwischen der Schicht, die die Titanverbindung mit kubischer Gitterstruktur enthält
20 und der Aluminiumoxidschicht oder zwischen den Aluminiumoxidschichten liegt, und
die Zwischenschicht Titanoxid mit einer Korund-Gitterstruktur und weiterhin Titancarbonitroxid mit einer kubi-
schen Gitterstruktur umfasst.

2. Gegenstand des Anspruchs 1, worin das Substrat Wolframcarbid umfasst.

25 3. Gegenstand des Anspruchs 1 oder 2, worin die mindestens eine Schicht, die die Titanverbindung mit kubischer
Gitterstruktur enthält, mindestens eine Schicht ausgewählt aus der Gruppe, bestehend aus Titancarbid, Titanitrid,
Titancarbonitrid, Titancarboxid, Titannitroxid und Titancarbonitroxid umfasst.

30 4. Gegenstand eines der Ansprüche 1 bis 3, worin die Zwischenschicht eine Dicke von 0,1 bis 5 μm hat.

5. Gegenstand eines der Ansprüche 1 bis 3, worin die Zwischenschicht eine Dicke von 0,05 bis 2 μm hat.

6. Gegenstand eines der Ansprüche 1 bis 5, worin die harte Deckschicht eine Dicke von 3 bis 25 μm hat.

35 7. Gegenstand eines der Ansprüche 1 bis 6, worin jede der Aluminiumoxidschichten eine Dicke von 0,5 bis 10 μm hat.

8. Gegenstand gemäß einem der Ansprüche 1 bis 7, worin die Zwischenschicht, die Titanoxid mit einer Korund-
Gitterstruktur enthält, eine maximale Peakintensität bei $2\theta = 34,5 \pm 1^\circ$ im Röntgenbeugungsmuster unter Verwen-
40 dung eines Cu $\text{k}\alpha$ -Strahls zeigt.

9. Gegenstand gemäß einem der Ansprüche 1 bis 8, worin das Atomverhältnis von Kohlenstoff, Stickstoff, Sauerstoff
und Titan in der Zwischenschicht wie folgt ausgedrückt wird:

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$$0 \% \leq (\text{C}+\text{N})/(\text{Ti}+\text{O}+\text{C}+\text{N}) \leq 10 \%$$

10. Gegenstand gemäß Anspruch 9, worin das Atomverhältnis

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$$0,5 \% \leq (\text{C}+\text{N})/(\text{Ti}+\text{O}+\text{C}+\text{N}) \leq 5 \%$$

ist.

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11. Beschichtetes Carbid-Schneidewerkzeugelement umfassend:

ein Wolframcarbid enthaltendes Substrat und
auf dem Substrat eine harte Deckschicht, die eine Dicke von 3 bis 25 μm hat,

worin die harte Deckschicht mindestens eine Schicht, die eine Titanverbindung mit kubischer Gitterstruktur enthält, mindestens zwei Schichten, die Aluminiumoxid enthalten und mindestens eine Zwischenschicht umfasst,

wobei die Zwischenschicht zwischen der Schicht, die die Titanverbindung mit kubischer Gitterstruktur enthält und der Aluminiumoxidschicht oder zwischen den Aluminiumoxidschichten liegt, und die Zwischenschicht Titanoxid mit Korund-Gitterstruktur und weiterhin Titancarbonitroxid mit einer kubischen Gitterstruktur umfasst.

12. Gegenstand des Anspruchs 11, worin die mindestens eine Schicht, die die Titanverbindung mit kubischer Gitterstruktur enthält, mindestens eine Schicht ausgewählt aus der Gruppe, bestehend aus Titancarbid, Titannitrid, Titancarbonitrid, Titancarboxid, Titannitroxid und Titancarbonitroxid umfasst.

13. Gegenstand gemäß Anspruch 11 oder 12, worin jede der Aluminiumoxidschichten eine Dicke von 0,5 bis 10 µm hat.

14. Gegenstand gemäß einem der Ansprüche 11 bis 13, worin die Zwischenschicht eine Dicke von 0,05 bis 2 µm hat.

15. Gegenstand gemäß einem der Ansprüche 11 bis 14, worin die Zwischenschicht, die Titanoxid mit einer Korund-Gitterstruktur enthält, eine maximale Peakintensität bei $2\theta = 34,5 \pm 1^\circ$ im Röntgenbeugungsmuster unter Verwendung eines Cu $\kappa\alpha$ -Strahls zeigt.

16. Gegenstand gemäß einem der Ansprüche 1 bis 15, worin die Zwischenschicht mit beiden, der Schicht, die die Titanverbindung mit kubischer Gitterstruktur enthält und mit der genannten Aluminiumoxidschicht, in Kontakt ist.

25 Revendications

1. Élément d'outil de coupe revêtu de carbure comprenant :

♦ un substrat ; et

♦ une couche de revêtement dure sur ledit substrat ;

➤ dans lequel ladite couche de revêtement dure comprend au moins une couche comprenant un composé du titane ayant une structure de réseau cubique, au moins une couche comprenant de l'oxyde d'aluminium, et au moins une couche intermédiaire ;

➤ dans lequel ladite couche intermédiaire est comprise entre ladite couche comprenant ledit composé du titane ayant une structure de réseau cubique et ladite couche d'oxyde d'aluminium, ou entre lesdites couches d'oxyde d'aluminium, et

➤ ladite couche intermédiaire comprend l'oxyde de titane ayant une structure de réseau du type corindon et comprend de plus le carbonitroxide de titane dans une structure de réseau cubique.

2. Article selon la revendication 1, dans lequel ledit substrat comprend du carbure de tungstène.

3. Article selon la revendication 1 ou 2, dans lequel ladite couche comprenant ledit composé du titane ayant une structure de réseau cubique comprend au moins une couche choisie dans le groupe formé par le carbure de titane, le nitride de titane, le carbonitrite de titane, le carboxyde de titane, le nitroxyde de titane et le carbonitroxide de titane.

4. Article selon l'une quelconque des revendications 1 à 3, dans lequel ladite couche intermédiaire possède une épaisseur de 0,1 à 5 µm.

5. Article selon l'une quelconque des revendications 1 à 3, dans lequel ladite couche intermédiaire possède une épaisseur de 0,05 à 2 µm.

6. Article selon l'une quelconque des revendications 1 à 5, dans lequel ladite couche de revêtement dure possède une épaisseur de 3 à 25 µm.

7. Article selon l'une quelconque des revendications 1 à 6, dans lequel chacune desdites couches d'oxyde d'aluminium possède une épaisseur de 0,5 à 10 µm.

8. Article selon l'une quelconque des revendications 1 à 7, dans lequel ladite couche intermédiaire comprenant de l'oxyde de titane ayant une structure de réseau du type corindon présente une intensité maximale à $2\theta = 34,5 \pm 1^\circ$ dans un diagramme de diffraction des rayons X utilisant la raie $\text{k}\alpha \text{ Cu}$.

9. Article selon l'une quelconque des revendications 1 à 8, dans lequel le rapport atomique de carbone, d'azote, d'oxygène et de titane dans ladite couche intermédiaire est exprimé comme suit :

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$$0\% \leq (\text{C}+\text{N})/(\text{Ti}+\text{O}+\text{C}+\text{N}) \leq 10\%$$

10. Article selon la revendication 9, dans lequel ledit rapport atomique est :

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$$0,5\% \leq (\text{C}+\text{N})/(\text{Ti}+\text{O}+\text{C}+\text{N}) \leq 5\%$$

11. Élément d'outil de coupe revêtu de carbure comprenant :

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- ♦ un substrat comprenant un carbure de tungstène ; et
- ♦ une couche de revêtement dure sur ledit substrat ayant une épaisseur de 3 à 25 µm ;

25 ➤ dans lequel ladite couche de revêtement dure comprend au moins une couche comprenant un composé du titane ayant une structure de réseau cubique, au moins deux couches comprenant de l'oxyde d'aluminium, et au moins une couche intermédiaire ;
 ➤ dans lequel ladite couche intermédiaire est comprise entre ladite couche comprenant ledit composé du titane ayant une structure de réseau cubique et ladite couche d'oxyde d'aluminium, ou entre lesdites couches d'oxyde d'aluminium, et
 ➤ ladite couche intermédiaire comprend l'oxyde de titane ayant une structure de réseau du type corindon et comprend de plus le carbonitroxyde de titane dans une structure de réseau cubique.

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12. Article selon la revendication 11, dans lequel ladite couche comprenant ledit composé du titane ayant une structure de réseau cubique comprend au moins une couche choisie dans le groupe formé par le carbure de titane, le nitrure de titane, le carbonitrure de titane, le carboxyde de titane, le nitroxyde de titane et le carbonitroxyde de titane.

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13. Article selon la revendication 11 ou 12, dans lequel chacune desdites couches d'oxyde d'aluminium possède une épaisseur de 0,5 à 10 µm.

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14. Article selon l'une quelconque des revendications 11 à 13, dans lequel ladite couche intermédiaire possède une épaisseur de 0,05 à 2 µm.

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15. Article selon l'une quelconque des revendications 11 à 14, dans lequel ladite couche intermédiaire comprenant de l'oxyde de titane ayant une structure de réseau du type corindon présente une intensité maximale à $2\theta = 34,5 \pm 1^\circ$ dans un diagramme de diffraction des rayons X utilisant la raie $\text{k}\alpha \text{ Cu}$.

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16. Article selon l'une quelconque des revendications 1 à 15, dans lequel ladite couche intermédiaire est en contact avec à la fois ladite couche comprenant ledit composé du titane ayant une structure de réseau cubique et ladite couche d'oxyde d'aluminium

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